



# THE UNIVERSITY *of* EDINBURGH

## Edinburgh Research Explorer

### **Novel combination of CRISPR-based gene drives eliminates resistance and localises spread**

**Citation for published version:**

Faber, N, McFarlane, G, Gaynor, C, Pocrnic, I, Whitelaw, B & Gorjanc, G 2021, 'Novel combination of CRISPR-based gene drives eliminates resistance and localises spread', *Scientific Reports*, vol. 11, 3719. <https://doi.org/10.1038/s41598-021-83239-4>

**Digital Object Identifier (DOI):**

[10.1038/s41598-021-83239-4](https://doi.org/10.1038/s41598-021-83239-4)

**Link:**

[Link to publication record in Edinburgh Research Explorer](#)

**Document Version:**

Publisher's PDF, also known as Version of record

**Published In:**

Scientific Reports

**Publisher Rights Statement:**

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. Te images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

**General rights**

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

**Take down policy**

The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact [openaccess@ed.ac.uk](mailto:openaccess@ed.ac.uk) providing details, and we will remove access to the work immediately and investigate your claim.





OPEN

# Novel combination of CRISPR-based gene drives eliminates resistance and localises spread

Nicky R. Faber<sup>1,2✉</sup>, Gus R. McFarlane<sup>3</sup>, R. Chris Gaynor<sup>4</sup>, Ivan Pocrnic<sup>1</sup>, C. Bruce A. Whitelaw<sup>3</sup> & Gregor Gorjanc<sup>1</sup>

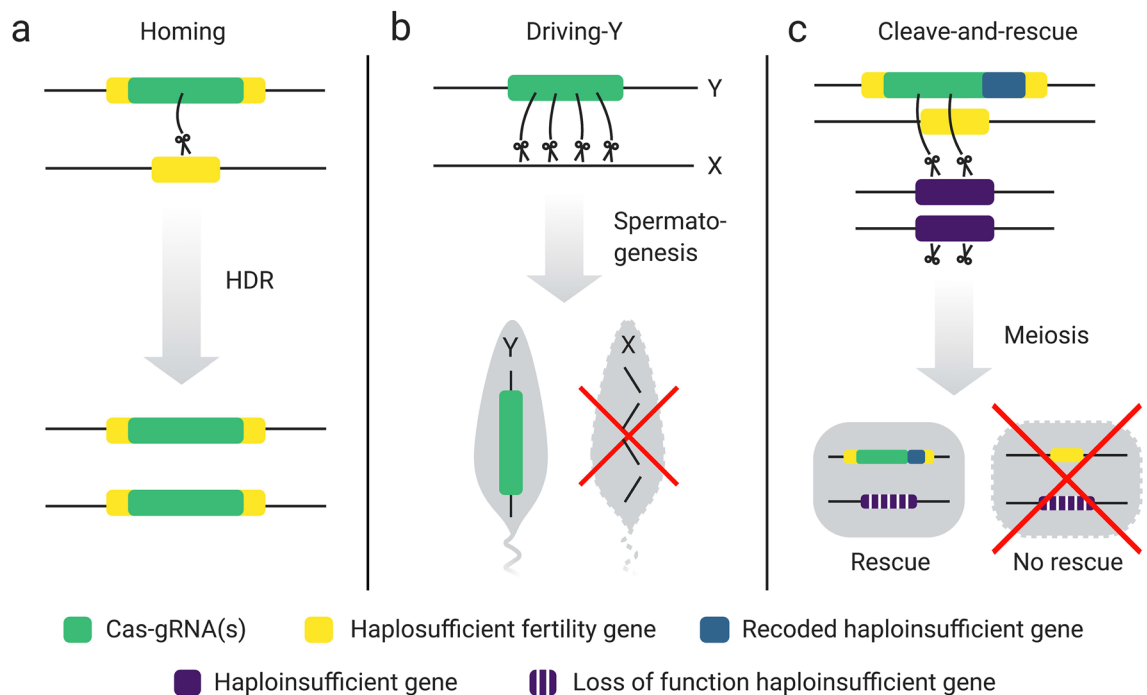
Invasive species are among the major driving forces behind biodiversity loss. Gene drive technology may offer a humane, efficient and cost-effective method of control. For safe and effective deployment it is vital that a gene drive is both self-limiting and can overcome evolutionary resistance. We present HD-ClvR in this modelling study, a novel combination of CRISPR-based gene drives that eliminates resistance and localises spread. As a case study, we model HD-ClvR in the grey squirrel (*Sciurus carolinensis*), which is an invasive pest in the UK and responsible for both biodiversity and economic losses. HD-ClvR combats resistance allele formation by combining a homing gene drive with a cleave-and-rescue gene drive. The inclusion of a self-limiting daisyfield gene drive allows for controllable localisation based on animal supplementation. We use both randomly mating and spatial models to simulate this strategy. Our findings show that HD-ClvR could effectively control a targeted grey squirrel population, with little risk to other populations. HD-ClvR offers an efficient, self-limiting and controllable gene drive for managing invasive pests.

CRISPR-based gene drives have the potential to address problems in public health, agriculture and conservation, including the control of invasive species<sup>1</sup>. Invasive species impact livelihoods, have severe economic consequences, and are among the major driving forces behind biodiversity loss<sup>2–4</sup>. Current control methods such as shooting, trapping, and poisoning are inhumane, labour-intensive, expensive, and ineffective in dealing with the scope of the problem in most situations<sup>5–7</sup>. Examples of damaging invasive species as a result of human mediated introduction include rabbits and cane toads in Australia, Asian carp in the US, and the grey squirrel and American mink in the UK.

In this study, we use the grey squirrel (*Sciurus carolinensis*) which is considered invasive in the UK as a case study for gene drive population control. First introduced in the 19th century, the grey squirrel is now widely distributed across the UK<sup>8</sup>. Since their introduction there has been a major decline in native red squirrels (*Sciurus vulgaris*). Grey squirrels are both larger and more aggressive than red squirrels and are passive carriers of Squirrelpox virus, which is lethal to red squirrels<sup>9</sup>. Without intervention, red squirrels could be lost from the UK mainland within the next few decades<sup>10</sup>. In addition to their impact on native red squirrels, grey squirrels also suppress natural forest regeneration through bark stripping of trees<sup>11</sup> and likely have a negative impact on biodiversity of native woodland birds by preying on eggs and chicks<sup>12</sup>. As an invasive pest they are estimated to cost the UK economy more than £14 million per year by debarking trees, gnawing through electricity cables and other forms of property damage<sup>13</sup>. A manageable and robust grey squirrel control strategy remains to be established<sup>7</sup>.

Although there are still technical challenges<sup>14,15</sup>, CRISPR-based gene drives may offer a humane, efficient, species-specific and cost-effective method for controlling invasive species, including grey squirrels in the UK<sup>16,17</sup>, filling a distinct void in the conservation toolbox. Broadly, a gene drive skews the inheritance ratio of an allele towards a super-Mendelian rate and therefore drives itself to spread quickly through a population<sup>18</sup>. The CRISPR-Cas system that these gene drives are based on comprises two components: a guide RNA (gRNA) and

<sup>1</sup>Highlander Lab, The Roslin Institute and Royal (Dick) School of Veterinary Studies, The University of Edinburgh, Easter Bush Campus, Midlothian EH25 9RG, UK. <sup>2</sup>Laboratory of Genetics, Wageningen University and Research, Droevendaalsesteeg 1, 6708PB Wageningen, The Netherlands. <sup>3</sup>Whitelaw Group, The Roslin Institute and Royal (Dick) School of Veterinary Studies, The University of Edinburgh, Easter Bush Campus, Midlothian EH25 9RG, UK. <sup>4</sup>AlphaGenes Group, The Roslin Institute and Royal (Dick) School of Veterinary Studies, The University of Edinburgh, Easter Bush Campus, Midlothian EH25 9RG, UK. ✉email: nfaber@outlook.com



**Figure 1.** Three CRISPR-based gene drives for population suppression. **(a)** Homing. A homing gene drive works by copying itself onto the homologous chromosome in the germline by directing Cas-gRNA(s) to cut a target site, which is repaired via homology directed repair (HDR). Therefore, all or most offspring inherit the gene drive. By locating Cas-gRNA(s) in the coding sequence of a haplosufficient female fertility gene, a female is fertile in homozygous state. All females are infertile once the gene drive allele is fixed leading to suppression of the population. **(b)** Driving-Y. During spermatogenesis, Cas-gRNA(s) are expressed from the Y-chromosome and shred the X-chromosome beyond repair. Therefore, all or most offspring from a driving-Y father will be driving-Y males. Population suppression is achieved by skewing the sex-ratio in favour of males. **(c)** Cleave-and-rescue. In the germline, Cas-gRNA(s) breaks an essential haploinsufficient gene whilst also supplying a recoded rescue version of this gene in the gene drive cassette. Therefore, only offspring which inherit the rescue within the gene drive are viable. Like the homing gene drive, the cleave-and-rescue gene drive can be located inside a haplosufficient female fertility gene, thereby making somatic homozygote females infertile and achieving population suppression.

a nonspecific Cas nuclease<sup>19</sup>. The gRNA directs the Cas nuclease to a specific sequence in the genome where it generates a double stranded break. Several synthetic CRISPR-based gene drives have been proposed with three major types suitable for population control: homing, driving-Y and cleave-and-rescue (Fig. 1)<sup>1,20–22</sup>. A homing gene drive works through a process called ‘homing’<sup>1</sup>. The system utilises germline-specific expression of CRISPR-Cas and subsequent cleavage in the germline, which leads to homology-directed repair (HDR) copying the gene drive element onto the homologous chromosome. By locating the homing gene drive cassette within the coding sequence of a haplosufficient female fertility gene, thereby disrupting the gene’s function, female somatic homozygotes will be infertile. As population growth is typically controlled by female reproductive performance<sup>18</sup>, the population will decline in size due to an increasing number of infertile females within the population. A driving-Y gene drive (also called X-shredder) specifically expresses CRISPR-Cas from the Y-chromosome during spermatogenesis to shred the X-chromosome at multiple locations beyond repair<sup>20,23</sup>. Therefore, only Y-bearing sperm mature and all or most offspring of a driving-Y father will inherit a gene drive harbouring Y-chromosome and be male. This eventually leads to a population decline due to the lack of breeding females. Cleave-and-rescue gene drive (also called toxin-antidote) uses CRISPR-Cas to cleave an essential gene while also supplying a recoded, uncleavable ‘rescue’ copy of this gene within the gene drive cassette<sup>21,24</sup>. In the case of a haploinsufficient essential gene, offspring must inherit the gene drive to be viable. For population suppression, the gene drive cassette is located in a haplosufficient female fertility gene like the homing gene drive, and thus this gene drive eventually also causes population decline due to female infertility<sup>24</sup>. In toxin-antidote terminology, this type of cleave-and-rescue gene drive is called Toxin-Antidote Dominant Embryo (TADE) suppression drive<sup>24</sup>.

Although all three population suppression gene drives are elegant and promising, they all face technical challenges. Homing gene drives face three major challenges. First, during in vivo testing, the formation of resistance alleles which block homing have been observed<sup>25,26</sup>. Resistance alleles can form through non-homologous end joining (NHEJ) instead of the desired homology-directed repair during homing. A potential solution is gRNA multiplexing<sup>16</sup>, but this is likely to reduce homing efficiency<sup>27,28</sup>. Second, another unresolved molecular challenge in mammals is the identification of a suitable promoter for sufficient, germline-specific expression of the gene drive. Ideally, the promoter should restrict Cas9 expression to a window when the endogenous meiotic

recombination machinery is most active<sup>15</sup>. Third, a homing gene drive that was not hindered by resistant alleles could theoretically spread indefinitely, thereby compromising global ecosystem safety. To address this concern, approaches to make gene drives self-limiting have been devised, including versions called ‘daisy drives’ (see Fig. 4A)<sup>29–32</sup>. Of these daisy drives the ‘daisyfield’ drive is the most straightforward mechanism to limit spread. In a daisyfield gene drive, the gRNAs are scattered throughout the genome (forming a daisyfield)<sup>32</sup>. These daisy elements are inherited in a Mendelian fashion, and therefore, offspring inherits half of the daisy elements from each parent. Thus, the gene drive stops spreading as the daisyfield is diluted through matings with wildtype individuals. Once all daisy elements have disappeared, all elements of the gene drive will likely also disappear due to negative selection (as homozygotes are infertile) and drift. This is desirable in case gene drive individuals spread to a non-target population. In a population where further spread is required, gene drive individuals with a complete daisyfield can be supplemented to keep the gene drive spreading. The rate and extent of suppression can be controlled by the number of gene drive animals supplemented and how many daisy elements the introduced animals carry. In contrast to homing gene drive, driving-Y gene drives face problems with the formation of a population equilibrium depending on shredding efficiency<sup>33,34</sup>. Furthermore, a major challenge in developing driving-Y in mammals is the identification of a highly-specific spermatogenesis promoter to drive Cas-gRNA expression<sup>17</sup>. Cleave-and-rescue gene drives have the advantage that multiplexing does not reduce efficiency as there is no homing involved, and therefore, the formation of resistance alleles is limited. Furthermore, cleave-and-rescue gene drives also show frequency-dependent dynamics, which can be exploited to keep the gene drive contained<sup>24</sup>. This poses practical challenges however, as it requires an accurate estimate of population size and the release of a large number of animals simultaneously.

Population control gene drives still need improvements for optimal efficiency, self-limitation, and controllability. In this study, we present HD-ClvR, a novel combination of gene drives that eliminates resistance, is self-limiting, and can be controlled in a practical manner. HD-ClvR is composed of homing (H), daisyfield (D), and cleave-and-rescue (ClvR) gene drives. Our modelling in grey squirrel demonstrates the strategy is highly efficient and overcomes the ongoing issue of resistance allele formation of homing gene drives. The daisyfield gene drive ensures self-limitation and allows for controlled, localised spread. Therefore, HD-ClvR could effectively control a targeted grey squirrel population, with little risk to other populations. Our analysis includes a randomly mating population and a spatially distributed population, which mimics the UK grey squirrel, though it can be adapted to other species. This study provides the first promising steps towards the development and testing of HD-ClvR.

## Results

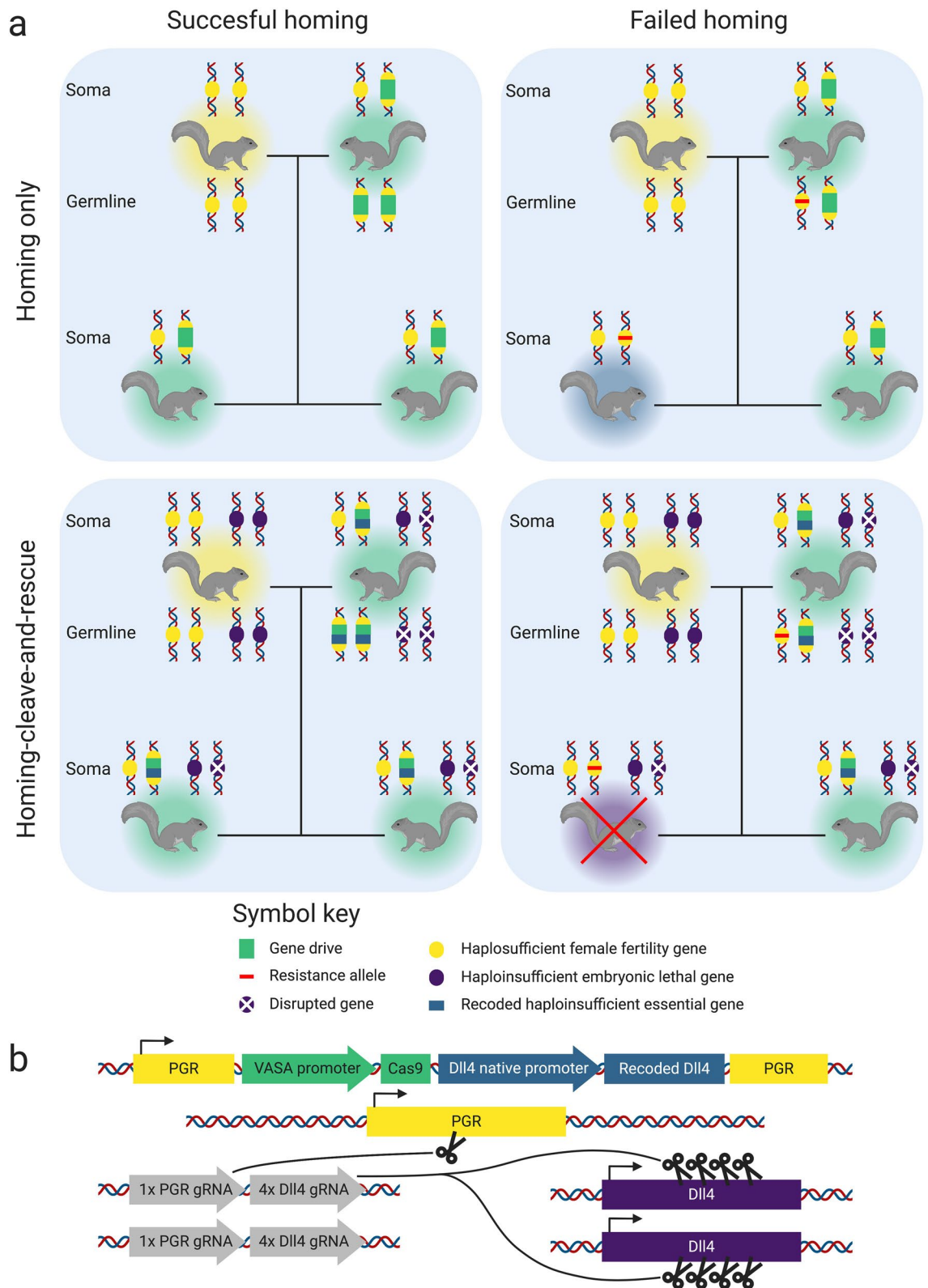
HD-ClvR is a combination of three gene drives: homing, daisyfield, and cleave-and-rescue. Our randomly mating and spatial modelling of this strategy in grey squirrel illustrates that HD-ClvR can effectively eliminate resistance allele formation, allows for optimised gRNA multiplexing, improves efficiency over standard cleave-and-rescue drives, and is both self-limiting and controllable. We find that the placement of supplemented animals significantly impacts the effectiveness of HD-ClvR, but that this is not prohibitive to the spread of the gene drive and that an effective placement strategy can achieve a rate of gene drive spread close to a randomly mating population.

**Eliminating resistance alleles.** By combining a homing gene drive with a cleave-and-rescue gene drive, HD-ClvR eliminates resistance alleles which occasionally form during gene drive homing (Fig. 2a). This works as follows: as germline homing occurs, both copies of a haploinsufficient essential gene are cleaved, and their function is destroyed through erroneous NHEJ-based repair. However, the homing construct contains a recoded, uncleavable copy of this haploinsufficient gene as a ‘rescue’. For offspring to be viable, they must inherit the gene drive with the rescue to have sufficient expression of the haploinsufficient gene. Offspring that inherit a resistance allele instead of the gene drive will not develop as they lack the rescue gene to compensate for their broken copy of the haploinsufficient gene. This mechanism prevents the emergence of resistance alleles.

HD-ClvR also allows for independent optimising of gRNA multiplexing for both homing efficiency and resistant allele elimination. Multiplexing gRNAs can overcome resistance allele formation, allowing homing to take place even if some resistant gRNA sites are present. With a standard homing gene drive, the optimal number of gRNAs is a trade-off between homing efficiency and overcoming resistance allele formation. Two gRNAs has been proposed as optimal for homing, with efficiency decreasing when more than two gRNAs are used<sup>28</sup>. However, to also limit the formation of resistance alleles, the optimal number in the trade-off lies between 4 and 8<sup>28</sup>. In contrast, with HD-ClvR it is possible to select the optimal number of gRNAs for homing, while multiplexing several gRNAs within the cleave-and-rescue to reduce the probability of resistance allele formation to effectively zero. Current data suggests four gRNAs is sufficient to prevent resistant allele formation<sup>28</sup>.

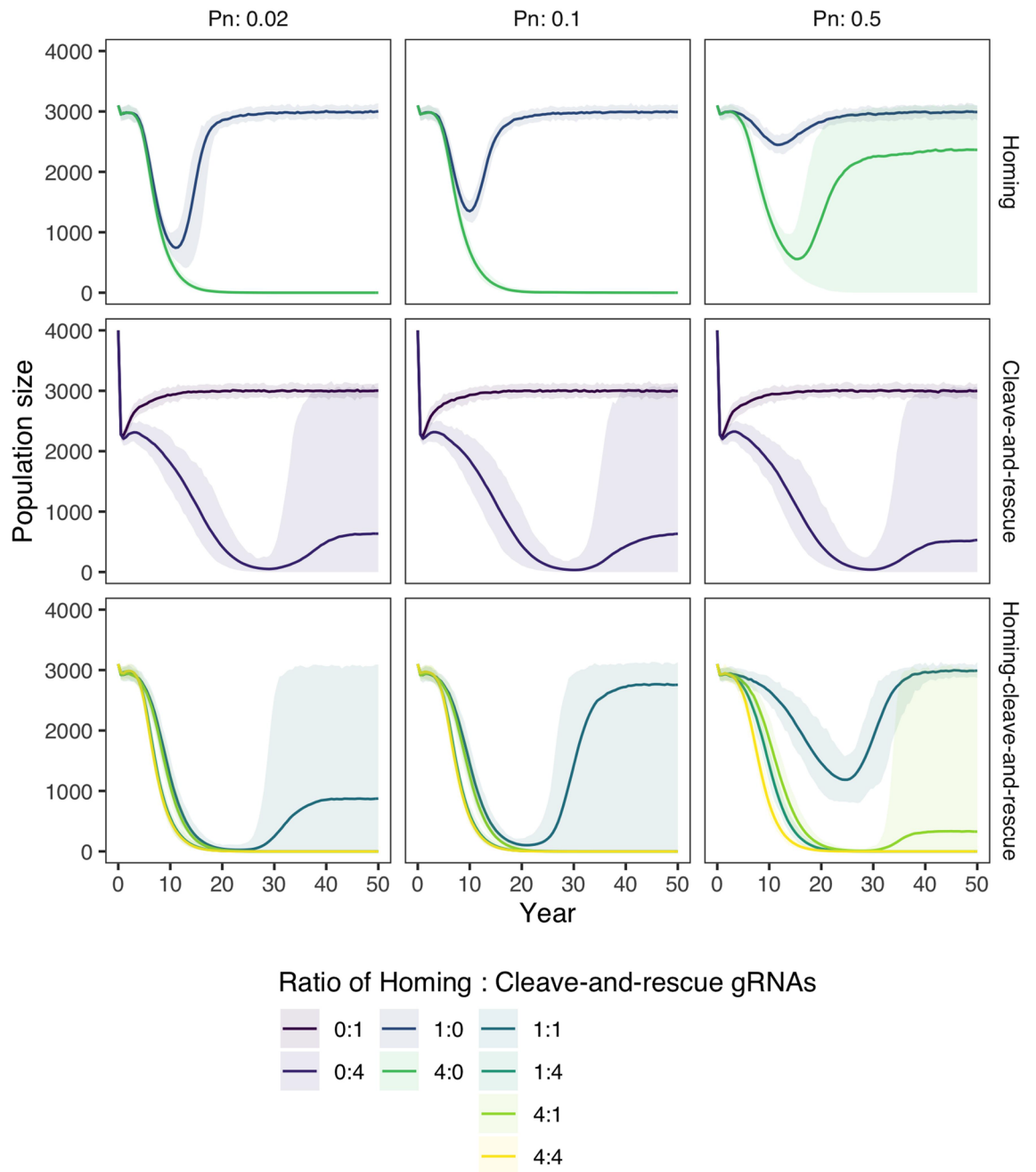
In grey squirrel, we have selected two genes through literature mining which are suitable for HD-ClvR: Progesterone Receptor (PGR) as a haploinsufficient female fertility gene involved in ovulation<sup>35</sup> and Delta-Like Canonical Notch Ligand 4 (DLL4) as a haploinsufficient essential gene involved in the Notch signalling pathway<sup>36</sup>. Both of these genes are conserved across many taxa (including vertebrates, arthropods, and nematodes among others) and if proven effective could also be used for other invasive species<sup>37</sup>. Figure 2b shows a candidate HD-ClvR construct design for grey squirrel control, using 1 gRNA for homing and 4 gRNAs for cleave-and-rescue.

To demonstrate that combining a homing and cleave-and-rescue gene drive can eliminate the formation of resistance alleles, we model a standard homing gene drive, a standard cleave-and-rescue gene drive, and a homing-cleave-and-rescue gene drive in a randomly mating population of grey squirrels. We model different rates of NHEJ ( $P_n$ , Fig. 3), which is the probability that a gamete is repaired through NHEJ instead of HDR. When NHEJ occurs, we model a probability of 0.33 that the target is a functional resistant allele, which is the probability that a frameshift did not occur. Like Prowse et al.<sup>16</sup>, we model no fitness cost to heterozygote gene drive animals. Our model uses either 1 or 4 gRNAs to show multiplexing reduces resistance allele formation. For



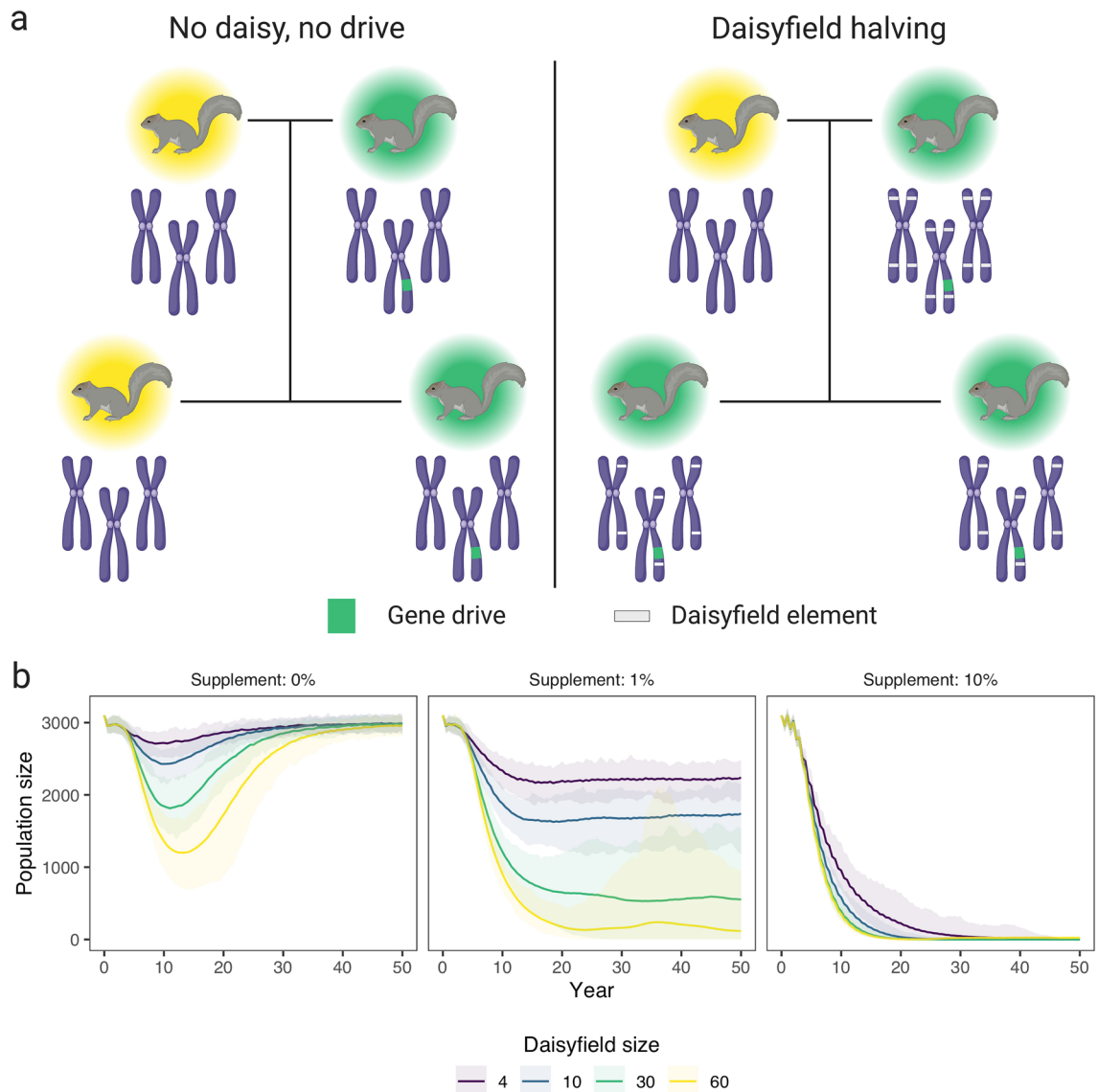
**Figure 2.** (a) A comparison of the inheritance scenarios of a homing-only gene drive (top row) and a homing-cleave-and-rescue gene drive (bottom row). The two panels in the left column show inheritance when homing is successful, and the two panels on the right show inheritance when homing fails. Each panel shows two parent squirrels and two offspring, each with the loci relevant for the gene drive. A legend for the gene drive components is provided. Squirrels colour coded halos represent their genotype: yellow = wildtype, turquoise = gene drive, blue = resistant, and purple = non-viable. (b) A potential HD-ClvR construct for grey squirrel. Colour coding is consistent with (a) and additionally, gRNAs are shown in grey. The gRNAs shown in this figure constitute one daisy element, multiple of these would constitute a daisyfield.





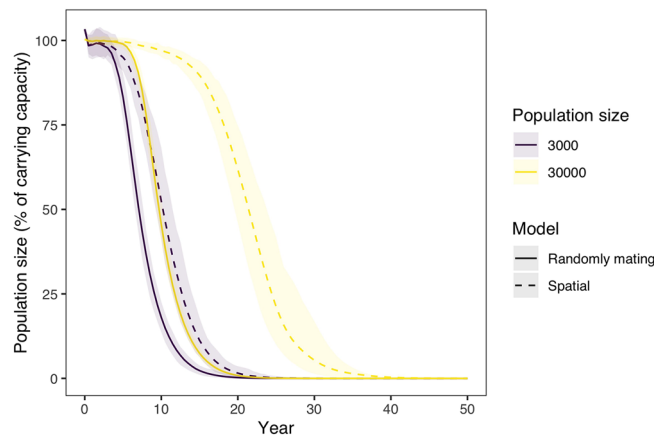
**Figure 3.** Population size over time after the introduction of gene drive squirrels with either a standard homing, a standard cleave-and-rescue, or a homing-cleave-and-rescue gene drive to a population with carrying capacity 3000. All simulations are based on a single release of 100 squirrels is done, other than the standard cleave-and-rescue gene drive, which requires a release of 1000 squirrels. Lines represent the average population size over 100 model replications, while opaque ribbons represent the 95% quantiles. The model was run with 3 different rates of NHEJ repair during homing ( $P_n$ ) and with different numbers of gRNAs for the homing and the cleave-and-rescue components of the gene drive.

the standard cleave-and-rescue gene drive, we modelled the release of 1000 gene drive squirrels instead of 100 gene drive squirrels, as this form of drive is only effective at a large introduction frequency. A standard homing gene drive was effective at low rates of NHEJ ( $P_n = 0.02$  and  $0.1$ ) when multiplexing 4 gRNAs but is inhibited by resistant alleles when only 1 gRNA is used at the same rates of NHEJ. However, at a higher rate of NHEJ ( $P_n = 0.5$ ), squirrels with resistant alleles rescue the population from standard homing gene drive suppression despite multiplexing 4 gRNAs. In contrast, with a homing-cleave-and-rescue gene drive, resistant alleles are eliminated, and the squirrel population is completely suppressed across all rates of NHEJ when 4 gRNAs are used in the cleave-and-rescue component of the drive. When we compare the three gene drive types in a large population of carrying capacity 30,000 instead of 3000, we see the same dynamics (Supplementary Figure S1).



**Figure 4.** (a) Two key aspects of a daisyfield gene drive: no daisy, no drive, and the halving of the daisyfield each mating with a wildtype individual. If there are no daisy elements in the genome, there is no drive due to the absence of a gRNA. Daisy elements are scattered throughout the genome (forming a daisyfield) and inherited in a Mendelian fashion. Therefore, offspring inherits half of the daisy elements from each parent. Thus, after enough matings with wildtype individuals, there will be no daisy elements left. (b) Population size over time after the introduction of 100 squirrels with a HD-ClvR gene drive to a population of carrying capacity 3000. Lines represent the average population size over 100 model replications, while opaque ribbons represent the 95% quantiles. The model was run with an NHEJ rate ( $P_n$ ) of 0.02, 1 homing gRNA, and 4 cleave-and-rescue gRNAs. Gene drive squirrel supplementation was done yearly, the amount being a percentage (0, 1, or 10%) of the total population size at that moment.

Although we model the homing gene drive component of HD-ClvR targeting a haplosufficient female fertility gene in this study, HD-ClvR is adaptable and could target any desirable gene to generate a loss of function mutation through insertion disruption or propagate a genetic cargo of interest. The cleave-and-rescue component of the HD-ClvR targets a haploinsufficient developmental gene in this study but this could also be adjusted to a haploinsufficient both-sex infertility gene. Our results suggest it is marginally more efficient to target an embryonic lethal gene (Supplementary Figure S2), as this prevents infertile resistant individuals from competing with gene drive individuals for resources. From an ethical standpoint the reduction in efficiency when targeting a both-sex fertility gene, instead of an embryonic lethal gene, may be justified by the improved societal and political acceptance for a strategy that evades killing and suppresses through infertility. Additionally, we tested if overexpression of the cleave-and-rescue target gene should be biologically tolerable, as individuals could have more than 2 copies of this gene if cleaving of the wildtype gene fails (Supplementary Figure S2). We conclude that when multiplexing sufficiently for the cleave-and-rescue part of the gene drive, there is no difference. As can



**Figure 5.** Population size over time after the introduction of 100 squirrels with a homing-cleave-and-rescue gene drive with 1 homing gRNA and 4 cleave-and-rescue gRNAs. The model was run for a randomly mating and a spatial model, and also for a small (carrying capacity 3000) and large population (carrying capacity 30,000). In the spatial model, gene drive squirrels were placed in the middle of the area. An NHEJ rate ( $P_n$ ) of 0.02 was used. Lines represent the average population size over 100 model replications, while opaque ribbons represent the 95% quantiles.

be seen from the dynamics when multiplexing less or not at all, allowing overexpression makes the gene drive initially faster to spread, but also allows resistance alleles to persist in the population.

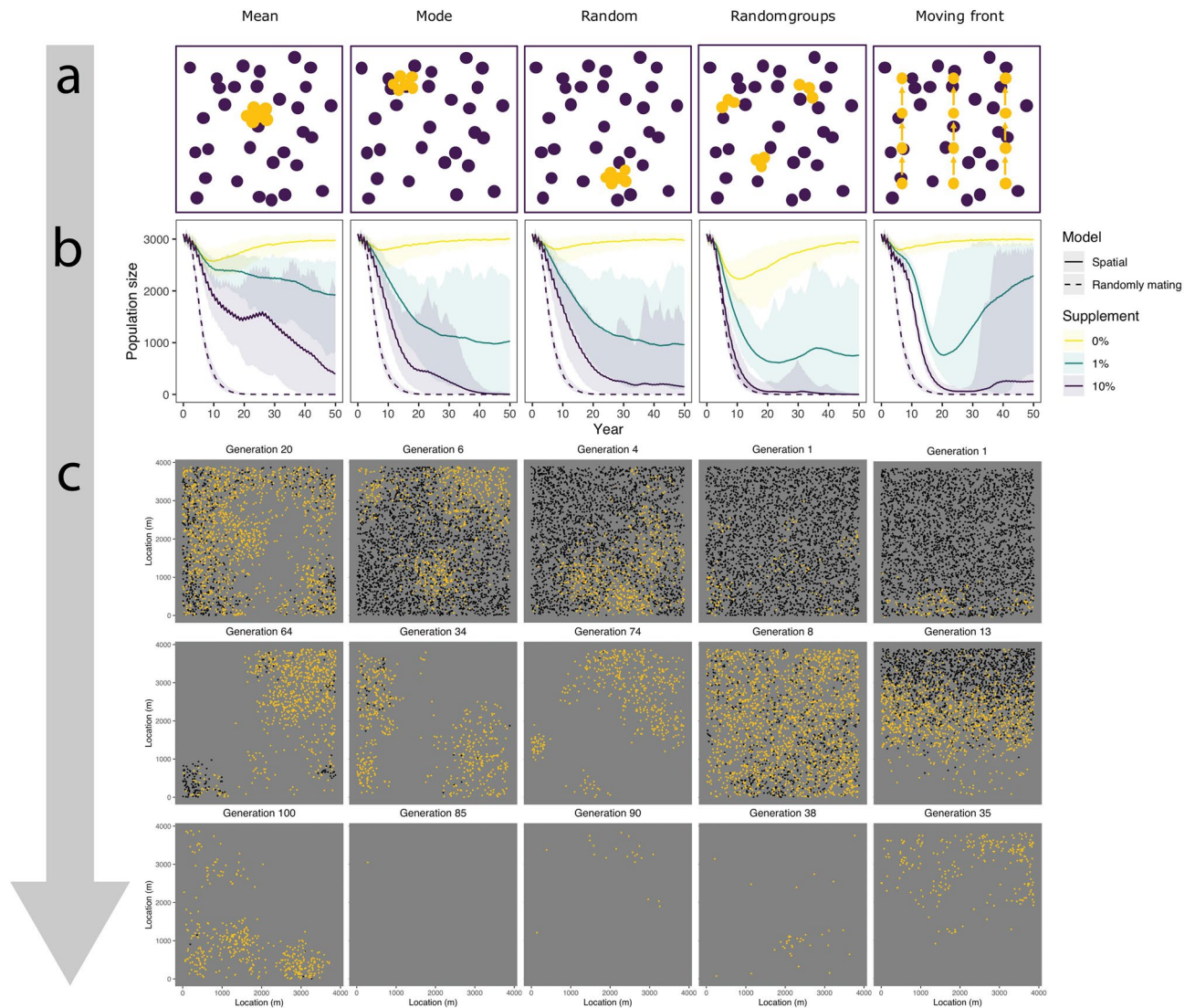
**Self-limitation and control.** A key benefit of HD-ClvR is that by including a daisyfield gene drive, it is self-limiting and can be controlled based on the number of supplemented gene drive animals and number of daisy elements each supplemented animal harbours (Fig. 4a). Unlike a standard homing gene drive, HD-ClvR can control the rate and extent of population suppression and, if required, suppression could be stopped by terminating further animal supplementation. Additionally, HD-ClvR does not require the large initial releases of standard cleave-and-rescue animals, which places pressure on the local ecosystem.

Using our randomly mating model, we show in Fig. 4b that by including a daisyfield system in a homing-cleave-and-rescue drive to form HD-ClvR, we can efficiently suppress a targeted population, while limiting risk to other populations, especially if those are bigger than the target population (Supplementary Figure S3). We modelled HD-ClvR with different daisyfield sizes in a population of 3000 grey squirrel over different rates of annual supplementation following an initial release of 100 HD-ClvR squirrels. The model shows that once the HD-ClvR runs out of daisy elements the population recovers. Therefore, HD-ClvR poses less risk to non-target populations than a standard homing gene drive. With 1% annual supplementation of HD-ClvR squirrels, the population size is reduced and maintained at an equilibrium, and with 10% annual supplementation the targeted population of grey squirrel is removed for all daisyfield sizes. Variability in the model with 1% supplementation and 60 daisyfield elements is caused by two factors: first, only a small number of individuals is supplemented due to the low population equilibrium, so randomness can have a big impact. Second, we round down the number of supplemented animals, to avoid perpetual maintenance of the population. Therefore with 1% supplementation, no animals are supplemented when the population is below 100 individuals. In Supplementary Figure S4, we show that it is possible to suppress a population without an accurate estimation of population size, which will be hard to obtain for most wild populations. To find the optimal combination of supplementation rate and daisyfield size, we ran a range of these two parameters and found that 5% supplementation would be sufficient to suppress a population, even with a small daisyfield (Supplementary Figure S5).

**Spatial dynamics and supplementation of HD-ClvR.** To understand the spatial dynamics of homing-cleave-and-rescue drives, initially excluding daisyfield, we modelled this approach in a simple spatial model. Modelling a single release of 100 homing-cleave-and-rescue gene drive squirrels in populations of 3000 and 30,000 squirrels, the model demonstrated that the spatial life history of grey squirrel allows for the spread of the gene drive (Fig. 5). We also show that the removal of the target squirrel population is more delayed in the spatial model than in the randomly mating population model. This difference is approximately 5 years in a small population, and is increased to approximately 15–20 years in a big population. To test the sensitivity of our model to two crucial parameters, mating range and migration range, we performed a sensitivity analysis and conclude that the model is sensitive to a decreased mating range, but not to a decreased migration range (Supplementary Figure S6).

Using our spatial model, we then explored how the placement of supplemented HD-ClvR animals could impact population suppression. We show the impact of different supplementation placement schemes by modelling five strategies: mean of population location, mode of population location, randomly, randomly in 10 groups, and in a moving front (Fig. 6a). The moving front was implemented such that we start at the bottom and move upwards in ten steps, thereafter, supplementing at the topmost location. As can be seen in Fig. 6b, different





**Figure 6.** Spatial dynamics of HD-ClvR using different placement schemes. **(a)** A schematic overview of the placement schemes. **(b)** Population size as a function of the placement schemes and amounts of supplementation. We modelled population size over time after the introduction of 100 squirrels with a HD-ClvR gene drive with 1 homing gRNA and 4 cleave-and-rescue gRNAs to a population of carrying capacity 3000. We modelled an NHEJ rate ( $P_n$ ) of 0.02 and a daisyfield of size 30. **(c)** Three snapshots of moments representing key spatial dynamics at 10% supplementation. See the full animations in Supplementary Videos S7.

placement schemes significantly affect the efficiency of the strategy. Placement at the mean population location was least effective and placement of squirrels randomly in 10 groups was most effective. Figure 6c shows three moments which represent key spatial dynamics of each placement scheme. For animations of the spatial dynamics over the whole timeline, see the animated GIFs (Supplementary Videos S7).

## Discussion

This research presents HD-ClvR, which is a combination of three gene drives: homing, cleave-and-rescue and daisyfield. Our modelling indicates that HD-ClvR overcomes an important trade-off in current homing gene drive designs: the trade-off between resistance allele formation and gene drive efficiency. This strategy benefits from the efficiency of a homing gene drive and the evolutionary stability of cleave-and-rescue gene drive. Due to the inclusion of a daisyfield system, HD-ClvR is self-limiting and can be controlled by supplementation of gene drive animals.

**HD-ClvR compared to other gene drives.** Over recent years, many different gene drives have been published and developments have been geared towards both efficiency and safety<sup>38</sup>. An ongoing issue has been the development of resistance alleles. For CRISPR-based homing gene drive there are two fundamental approaches to combat resistance allele formation: careful gRNA targeting and gRNA multiplexing. When a gRNA targets a conserved sequence in a gene, resistance alleles are likely to disrupt gene function through NHEJ repair and

will therefore reduce fitness<sup>39</sup>. Recently, population suppression was already shown to work with a carefully targeted homing gene drive in contained mosquito populations<sup>39</sup>, however, current data suggests that homing might be less efficient in mammals than in insects<sup>14</sup>. A recent paper has proposed the concept of 'tethered homing gene drive', which combines a threshold-dependent underdominance gene drive with a homing gene drive for improved suppression capabilities<sup>40</sup>. We use this concept in a different manner in HD-ClvR, by relying on a daisyfield rather than threshold-dependence for self-limitation. Very recently, two new papers have proposed a gene drive similar to HD-ClvR, but intended for population modification instead of suppression<sup>41,42</sup>. These studies also combine homing and cleave-and-rescue principles to combat resistance alleles and their modifications are able to persist stably in cage experiments, which is promising for HD-ClvR.

In addition to targeting conserved sequences, when gRNA multiplexing, resistant allele formation is reduced because multiple sites are targeted simultaneously. For homing gene drives, multiplexing has been shown to reduce homing efficiency when more than two gRNAs are used<sup>28</sup>. In contrast, cleave-and-rescue gene drives do not have this problem, as they do not use homing and can therefore multiplex gRNAs without any efficiency costs. HD-ClvR separates the elimination of resistance alleles and homing efficiency, and therefore gRNAs can be optimised for both goals separately.

To date, most gene drive research has focused on improving the efficiency, however, equally important is the development of strategies that allow for containment, or even reversibility, of the gene drives<sup>29,43</sup>. For contained gene drives, density dependence is often used, which requires large numbers of gene drive individuals to be released into a target population to spread<sup>44</sup>. Therefore, non-target populations are unlikely to be affected by this type of gene drive. However, a large single release of gene drive individuals can put significant pressure on the local ecosystem, and if a population is already at carrying capacity, it may lead to starvation or mass migration of the population. In contrast, HD-ClvR uses ongoing input in the form of gene drive animals to control the extent of population suppression and contain spread, while the total amount of gene drive animals necessary for release is similar to threshold-dependent gene drives. Therefore, the use of HD-ClvR seems more feasible than threshold-dependent gene drives. Although self-limitation comes with increased cost and labour relative to unlimited gene drives, we believe this is justified by the control and safety of HD-ClvR.

As stated above, the initial introduction frequency for a standard cleave-and-rescue gene drive in our randomly mating model was increased 10-fold over the other homing-based strategies. This increase is necessary due to the significant cost to the reproduction rate that is incurred when using a standard cleave-and-rescue gene drive. On average, cleave-and-rescue animals will produce 50% less offspring than wild-type animals<sup>21,24</sup>. This significantly slows the spread of the gene drive and due to density dependent dynamics, requires large initial releases of cleave-and-rescue animals for population suppression. With a homing-cleave-and-rescue drive, more offspring inherit the drive and there is less cost to the reproduction rate. Effectively, for homing-cleave-and-rescue, the reproduction rate of gene drive individuals is equal to the homing efficiency (plus half of the homing failure rate, where the gene drive is inherited by chance), which so far has been shown to range from 0.7 to 1 in different organisms<sup>14,39,45</sup>.

**Supplementation.** As animal supplementation is a critical component of HD-ClvR, our modelling investigated how daisyfield size and the level and placement of supplemented HD-ClvR animals effects efficiency and safety of population suppression. Optimisation of these parameters can significantly reduce cost and labour, as well as reduce the risk of unwanted impacts on non-target populations. We modelled our supplementation as a percentage of the total population size, therefore the number of individuals needed for supplementation increases linearly with population size. We also want to minimise the risk of non-target populations being impacted by the gene drive, and therefore, there is a trade-off between safety (size of the daisyfield) and cost and labour (level of supplementation required).

The least number of daisy elements that can suppress the population with a realistic level of supplementation, but does not cause any serious issues in non-target populations, should be objectively established through an in-depth risk assessment process. In a larger population however, the spread is slower than in a small one. Therefore, for improved safety and efficiency, gene drives are best applied in small sub-populations separately. The impact of a single introduction, such as a rogue deployment or migration, depends on the population size. The smaller the population, the bigger the impact. This is a concern when the target population is much larger than the non-target population, but this is not the case for invasive UK grey squirrels and many other invasive species.

The appropriate daisyfield size also depends on the rate of NHEJ ( $P_n$ ) of the gene drive system; the higher the ( $P_n$ ), the more embryonic lethal offspring will arise and the sooner daisyfield burns out. To choose a safe number of daisy elements, we also need an estimate of how many animals a rogue party could obtain, potential breed and add into a non-target population for their own benefit. Overall, each target population and prospective gene drive strategy needs to be considered on a case-by-case basis and include an in-depth multidisciplinary risk assessment process.

When we consider the spatial aspects of a HD-ClvR supplementation programme, the picture becomes more complex. A key factor is the supplementation location of individuals. Obviously, supplementing individuals in a location where the population has already been suppressed will be ineffective. Therefore, different placement strategies can be adopted to keep placing individuals in a relevant area. A monitoring system where not only the size of the population is known, but also the location can significantly help HD-ClvR continue spreading and suppress a targeted population.

In this study, we modelled HD-ClvR using five different supplementation placement strategies in grey squirrel. These were: supplementation at the mean of population location, the mode of population location, randomly, randomly in 10 groups, and in a moving front (Fig. 6a). With supplementation at the mean of the population location, supplementation started in the middle of the population. After a few generations, a gap appears in the

middle due to local suppression. The mean of the populations location still lies in the middle, as can be seen in Fig. 6c at 20 generations. Therefore, supplementation is not effective until the population is also suppressed in another location, thereby shifting the mean. Additionally, when there is a single large patch of the population left and additional smaller clusters, supplementation in the middle of the large patch allows the smaller clusters to recover, as can be seen in Fig. 6c after 64 generations.

With supplementation at the mode of the population location, we supplement in a location where there are many individuals. This placement strategy avoids the problem of supplementing in a location without individuals, either in a doughnut-like spatial population structure or in a multi-patch population. However, this placement strategy still allows small patches to form and recover. Supplementation at a random location theoretically means that supplementation happens uniformly, but in reality, this is not the case. Initially HD-ClvR spreads in multiple locations, but after the population is suppressed in certain regions, supplementation in those regions becomes ineffective. Therefore, at a later stage of population suppression this placement scheme becomes increasingly ineffective.

Supplementation at random locations is more effective when they are broken up into multiple groups (ten in our model). The gene drive spreads in many locations initially like the random single location placement scheme. After significant suppression of the population some but not all of the 10 groups supplemented are at ineffective locations. The groups that are placed at relevant locations are enough to keep the gene drive spreading. In our model supplementation in groups at random locations gets close to the speed at which a gene drive spreads in a non-spatial model.

The moving front placement scheme is very effective initially, as the gene drive spreads uniformly across the front. In this case, supplementation keeps ahead of where the populations is being suppressed. This placement strategy allows the population to recover behind the moving front after effective initial spread and near-complete suppression. To improve efficiency of the moving front strategy, it may be beneficial to include random supplementation behind the moving front to prevent animals from re-establishing.

Finally, in our spatial model, it was evident that there is more uncertainty in levels of population suppression than a randomly mating model leads us to believe. As can be seen in Fig. 6b, the 95% quantiles are broader than the quantiles in Fig. 3. Therefore, we conclude that to tailor the amount of supplementation, it is vital to closely monitor a population where a gene drive is used.

**Assumptions and future work.** Our model works under the following six assumptions. First, our model excludes some complexities of the optimal number of gRNAs for homing. Although our model suggests that multiplexing gRNAs for both the homing and cleave-and-rescue gene drives is most effective, a recent study using a more complex model and in vivo data shows that the optimal number of gRNAs to use for homing in *Drosophila melanogaster* is two. They report a decrease in homing efficiency with more than two gRNAs due to reduced homology and Cas nuclease saturation<sup>28</sup>. Therefore, our gene drive with four gRNAs for both homing and cleave-and-rescue will likely be less efficient in such a complex model. We suggest using two homing gRNAs and four cleave- and-rescue gRNAs is likely most efficient, while still eliminating all resistance alleles<sup>28</sup>. It would be prudent to analyse our gene drive in this complex model as well to get a definitive estimate, as Cas saturation is thought to have an influence on gene drive efficiency when multiplexing is used<sup>28</sup>.

Second, we assumed there was no embryonic Cas-gRNA expression. Embryonic Cas-gRNA expression might be problematic as it leads to resistance allele formation and can interfere with the cleave-and-rescue mechanism by cleaving alleles from the wildtype parent. As our gene drive eliminates resistance alleles, embryonic Cas-gRNA expression may not inhibit spread, depending on the rate. Additionally, if the embryonic Cas-gRNA expression turns out to be more common in grey squirrel or other species, the cleave-and-rescue part of the gene drive can be harnessed with a double rescue mechanism to overcome this issue, as reported by Champer et al.<sup>24</sup>.

Third, we did not take other types of resistance alleles into account such as mutations rendering the CRISPR-Cas non-functional. As this is a universal assumption in gene drive research, we will have to await multigenerational studies to see if this is problematic.

Fourth, HD-ClvR has not been tested in vivo, which is our next step. The two recent papers testing a gene drive similar to HD-ClvR for population modification have performed in vivo tests in *Drosophila melanogaster* which showed very efficient conversion rates<sup>41</sup>. Proof-of-concept testing of HD-ClvR would likely initially occur in *D. melanogaster* and mouse models before progressing to squirrel studies. Recent reports have shown that the VASA promoter for Cas expression in homing gene drives is not optimal and further investigation to identify a meiosis-specific germline promoter is needed<sup>15</sup>. Furthermore, the integration of many daisies in a squirrel genome will be a molecular challenge and is a feat which has not yet been reported on in any species. This task could be achieved using either a random integration strategy, such as lentiviruses<sup>46</sup> or a targeted integration strategy that exploits neutral repetitive sequences in the genome as target sites<sup>32</sup>. Also, non-model species might be difficult to genetically engineer, although grey squirrel embryology will likely follow the extensive knowledge on rodent and farmed animal embryology, and similar reagents and equipment could be used. An important consideration when engineering gene drive is that the modified animals maintain enough wild vigour to survive and breed in a wild population. Promising technologies for generating gene drive harbouring mammals with as little intervention as possible include in situ delivery of CRISPR reagents to the oviduct<sup>47</sup>.

Fifth, for our spatial modelling, we assumed that an estimation of population size could be made every year, although there is a significant amount of room for error in this estimate. Additionally, for some of our placement schemes, we assumed an accurate estimate of population location. As the random placement in groups scheme turned out most effective, this is not a problem so much as further potential for improvement. Another direction for future spatial work is the modelling of real landscapes, which are more complex than what we modelled in this study<sup>48</sup>. In complex landscapes, it might be that gene drive spread is slower or even regionally confined in

Parameter	Value	Source
<b>Population and reproduction</b>		
Population carrying capacity	3000	<sup>50</sup>
Maximum population growth rate*	1.16	<sup>51</sup>
Average litter size	2.87	<sup>52</sup>
Generation time (weeks)	26	<sup>53</sup>
<b>Spatial distribution</b>		
Home range radius (m)	80	<sup>54</sup>
Maximum density (individuals/home range)	4	<sup>54</sup>
Maximum mating range radius (m)	600	<sup>55</sup>
Mating range observations	30	<sup>55</sup>
Maximum dispersal range radius (m)	10000	<sup>56,57</sup>

**Table 1.** Key parameters used in the model for the grey squirrel. For the rest of the parameters, see the supplementary code. \*Calculated as the  $\log(\max\{R_0\})$ .

some situations. Additionally, there might be spatial dynamics to gene drives in general such as 'chasing', which is the perpetual escaping and chasing of wildtype and gene drive animals<sup>34</sup>. Further efforts are necessary to create a more realistic spatial model before we can consider using a gene drive.

A final consideration is that the ecological services the grey squirrel and other invasive species provide are largely uncharted. Ecologists need to investigate the ecological services that an invasive species performs and how an abrupt suppression of this invasive population might impact the ecosystem as a whole. We need to consider other restorative measures such as reintroducing native species to fragmented habitats, amongst other ecological interventions<sup>49</sup>. From a regulatory perspective, there is no tested legislative framework for the release of gene drive organisms; and with regard to our test animal it is currently illegal to breed grey squirrels in the UK. Developing these legislative frameworks alongside gene drive research is important. More importantly, the UK needs to continue to broaden public engagement and see whether the public is receptive to the deployment of gene drive technology in parallel to a financial overview of how much it would cost to apply gene drives reflecting our predicted need for supplementation.

**Summary.** HD-ClvR offers an efficient, self-limiting, and controllable gene drive strategy. We show that in the spatial model, complete population suppression is achieved approximately 5 years later than in the randomly mating population model. We then explored how the placement of supplemented animals could impact population suppression. Our results show that spatial dynamics of supplementation placement are not prohibitive to the spread of the gene drive, but that in fact, with an optimised strategy, spread at a rate equal to randomly mating population can be achieved. In our models, we have shown that grey squirrels have a spatial life history which facilitates the spread of a gene drive. Therefore, gene drives could be a valuable tool in the conservation toolbox.

## Methods

We describe our methods and materials in two sections. The first section details the randomly mating population model, and the second the spatial model. For the modelling, we adopted the work of Prowse et al.<sup>16</sup> and implemented new features. This model is an individual-based, stochastic, discrete-time model of a randomly mating population. Per individual, the model keeps track of several characteristics such as age, sex, parents, and the state of genetic loci involved in the gene drive. For each offspring, we model the homing and subsequent inheritance of the gene drive. By running this stochastic model several times, we obtain an impression of the possible outcomes. Several life history parameters of an organism are needed to run this model. The parameters we used to model a grey squirrel population can be seen in Table 1.

**Randomly mating model.** For the randomly mating model, we added three additional features to the model of Prowse et al.<sup>16</sup>: cleave-and-rescue, daisyfield, and driving-Y. Cleave-and-rescue and daisyfield were not tested by Prowse et al.<sup>16</sup>, who only compared homing-based gene drives. We also modelled a driving-Y-cleave-and-rescue gene drive, but the homing-cleave-and-rescue was deemed more promising because the identification of a highly-specific spermatogenesis promoter remains a challenge. In addition to these three new features, we extended the supplementation functionality, because daisyfield-based population suppression requires flexible supplementation.

1. *Cleave-and-rescue.* In the model, we keep track of each gRNA-targeted site in cleave-and-rescue target genes and their functionality in each individual. The homing gene drive construct contains the recoded rescue copy of this target gene. All wildtype organisms start with two viable target genes, while gene drive organisms start with one viable target gene and one rescue. In general, after germline Cas-gRNA activity, viable target genes are cleaved and the rescue gene homes along with the gene drive. However, as with any sites targeted by a Cas-gRNA, it is possible that resistance alleles form after non-homologous end joining and on occasion restore functionality of the target gene. Therefore, we implemented cleave-and-rescue gRNA multiplexing



in the model. The probability that cleave-and-rescue target genes go from  $i$  to  $j$  functional cutting sites ( $P_{ij}$ ) is:

$$P_{ij} = \binom{i}{i-j} (P_c(1 - P_f))^{i-j} (1 - P_c)^j P_b^{i-j-1}, \quad (1)$$

where  $P_c$  is the probability of cutting at a gRNA-targeted site,  $P_f$  is the probability of functional restoration in case of cutting, and  $P_b$  is the probability that a block of DNA in between two cutting sites is not removed. This formula consists of four factors: first, we multiply by all permutations of cutting sites, because their order is irrelevant. Second, we multiply by the probability that  $i - j$  cutting sites are all cut and repaired functionally. Third, we multiply by the probability that  $j$  sites remain uncut. Fourth, we multiply by the probability that no blocks of DNA in between cut sites were removed. We use  $P_c = 0.95$  and  $P_f = 0.667$  following Prowse et al.<sup>16</sup>. We estimated  $P_b$  from our unpublished data of 18 mouse embryonic stem cell lines, each cut simultaneously with Cas9 at two sites spaced 36 bp apart. In 3 out of 18 cases, the block of DNA in between the cut sites was not removed and therefore, we use a  $P_b$  of 0.2. All left-over probability ( $1 - P_{ij}$ ) is the probability that a target gene is rendered non-functional. An organism needs to have exactly two copies of the target gene (recoded rescue or original) to be viable. We assumed that there is no embryonic Cas-gRNA activity. After random inheritance of parental alleles, we remove non-viable offspring.

2. **Daisyfield.** We implemented daisyfield by tracking the number of daisyfield elements in the genome of each individual. Wildtype organisms start without any daisy elements and the number of daisy elements for gene drive organisms is a parameter in the model. Each daisy element contains both the homing and the cleave-and-rescue gRNAs and in case of gRNA multiplexing, it contains one of each different gRNA. Therefore, during germline Cas-gRNA expression, if no daisy elements are present, both homing and cleave-and-rescue can not occur. We assumed that daisy elements remain complete through every meiosis, so there is no crossing over in the middle of them. Also, we assumed that there is no linkage between daisy elements, that is, they are spaced far apart or located on different chromosomes. During inheritance, each daisy element from the parents has a 0.5 probability of being inherited to the offspring.
3. **Driving-Y.** Although the driving-Y is not a part of our final gene drive strategy, we implemented it in the model. The driving-Y gene drive is modelled on the Y-chromosome and skews the sex ratio of offspring towards males. The efficiency of this skew is a parameter in the model and is defined as the probability that offspring of a gene drive animal is male.
4. **Supplementation.** We made two changes to the supplementation already implemented by Prowse et al.<sup>16</sup>. Instead of yearly supplementation of the same amount as the initial gene drive release, we added two parameters to vary supplementation amount and interval. Supplementation amount can be any percentage of the total population size, and supplementation interval can be any decimal number of years as long as they coincide with generations.

**Spatial modelling.** For the spatial modelling, we added basic spatial functionality on top of the other additions to the randomly mating model of Prowse et al.<sup>16</sup>. We model a square, two-dimensional space and assume uniformly distributed resources such as food. The spatial functionality is comprised of four steps: spatial setup, distance-dependent mate allocation, offspring placement, and movement. The spatial setup is only done once at the start of the model and initiates everything necessary for spatial functionality. Mate allocation, offspring placement, and movement occur each generation, and their purpose is to reflect spatial life histories. Distance-dependent mate allocation ensures that squirrels who are close together are more likely to mate than squirrels further apart. Offspring placement demonstrates the location of birth and maternal care of individuals. Movement reflects the migration of individuals whenever overpopulation occurs in an area. With several parameters shown in Table 1, this spatial functionality can be adapted to reflect the spatial life history of many species. Additionally, we have added spatial placement strategies for supplementation.

1. **Spatial setup.** The first step in spatial modelling is to determine the size of the area in which the simulations take place. As we use a square two-dimensional space, we need to know the length of the side of this area  $A$ . We calculate  $A$  using the carrying capacity of the population  $K$ , the radius of the home range of the organism  $r$ , and the density at carrying capacity  $D$ :

$$A = \frac{\sqrt{K\pi}r^2}{\sqrt{D}}. \quad (2)$$

Essentially, this formula transforms a circular home range radius into an area, multiplies it by the number of individuals, transforms it into the length of a square area, and makes it smaller according to the density at carrying capacity. Using this formula, the area is exactly large enough to hold  $K$  number of individuals at  $D$  density. In this two-dimensional area, we track the  $x$  and  $y$  coordinates of individuals. Each individual starts at a random location within the area. Where gene drive individuals are placed depends on the placement strategy.

2. **Distance-dependent mate allocation.** During the reproduction step of the model, instead of random mate allocation, we use distance-dependent mate allocation. We do this in three steps. First, we calculate the Euclidian distance between all females and males. Second, we use a Gaussian radial basis function to calculate the probability of a male approaching the female to mate ( $P_a$ ), depending on the distance  $s$  between them:



$$P_a = e^{-(\epsilon s)^2}, \quad (3)$$

where the value  $\epsilon$  determines the shape of the radial basis function and is calculated from the mating range parameter. In the case of the grey squirrel, the maximum observed mating range was 600 out of 30 observations<sup>55</sup>. Therefore, we assumed that the probability of a mating range of 600 was 1/30 and from this, we calculate  $\epsilon$ . Third, from the males that do approach the female, we choose a random one as the father of the offspring. In the case that no males approach the female, she does not reproduce.

3. *Offspring placement.* We place offspring at the location of the female at the moment of reproduction.
4. *Movement.* In grey squirrels, migration is the driving force behind a stable population size<sup>58</sup>. Therefore, we implemented density-dependent migration and not density-dependent mortality. In the model, we make a distinction between the movement of migrants and residents. Firstly, we determine which individuals migrate and which remain as residents. This distinction is density dependent, that is, the density at the location of an individual determines the probability that they migrate ( $P_m$ ):

$$P_m = \begin{cases} 0 & d \leq D \\ 1 - \frac{D}{d} & d > D \end{cases}, \quad (4)$$

where the local density  $d$  and the density at carrying capacity  $D$  are measures of the number of individuals that are in the home range of an individual. Therefore, when the local density is below maximum density, individuals will not migrate. When the local density is higher than the maximum density, the probability of migration is equal to the proportion of individuals that need to migrate to leave the local density at the maximum density. Next, for both the resident and the migrant movement, we choose a direction and a distance to determine a new location. We choose a random direction and a distance from two separate gamma distributions for residents and emigrants with shape and scale parameters:  $distance \sim \Gamma(k, \theta) \equiv \text{Gamma}(5, r/5)$  for residents and  $distance \sim \Gamma(k, \theta) \equiv \text{Gamma}(5, 3r/5)$  for migrants,  $r$  being the home range. We use a broader distribution for migrants than for residents as migrants tend to travel greater distances<sup>55</sup>. The residents move to a random location in a single step. If the new location is out of the boundaries of the spatial space, we pick a new direction and distance. In contrast, migrants move in multiple steps within a certain migrational range to a place where there is space available, that is, where the local density  $d$  is lower than the density at carrying capacity  $D$ . The migrant searches for a new location in a lazy manner, which means that an animal will first try nearby locations, and incrementally migrate further if necessary. In each step, we pick a random direction and add a new distance from the gamma distribution to the previous distance. If the maximum migration distance is surpassed, the distance is set to zero and the process starts again. To ease the computational burden of this algorithm, we limit the number of steps to 50 and then, we keep the last location regardless of density.

5. *Supplementation.* The placement of individuals for supplementation is important. Therefore, we have implemented five placement strategies that can be used, although further exploration of this aspect is interesting. The six placement strategies are: middle of the area, mean of population location, mode of population location, random location at each supplementation, divided into 10 groups and placed at random locations at each supplementation, and divided into 10 groups and placed as a moving front in 10 steps.

## Code availability

Code is available from the Highlanderlab gitlab: [https://git.ecdf.ed.ac.uk/HighlanderLab\\_public/nfaber\\_squirrel\\_gd](https://git.ecdf.ed.ac.uk/HighlanderLab_public/nfaber_squirrel_gd).

Received: 6 November 2020; Accepted: 27 January 2021

Published online: 04 March 2021

## References

1. Esvelt, K. M., Smidler, A. L., Catteruccia, F. & Church, G. M. Emerging technology: Concerning RNA-guided gene drives for the alteration of wild populations. *Elife* **3**, e03401. <https://doi.org/10.7554/eLife.03401> (2014).
2. Mooney, H. A. Invasive alien species: The nature of the problem. *Scope Sci. Committee Probl. Environ. Int. Council Sci. Unions* **63**, 1 (2005).
3. Pejchar, L. & Mooney, H. A. Invasive species, ecosystem services and human well-being. *Trends Ecol. Evol.* **24**, 497–504. <https://doi.org/10.1016/j.tree.2009.03.016> (2009).
4. Sala, O. E. *et al.* Global biodiversity scenarios for the year 2100. *Science* **287**, 1770–1774. <https://doi.org/10.1126/science.287.5459.1770> (2000).
5. Luque, G. M. *et al.* The 100th of the world's worst invasive alien species. *Biol. Invas.* **16**, 981–985. <https://doi.org/10.1007/s10530-013-0561-5> (2014).
6. Campbell, K. J. *et al.* The next generation of rodent eradications: Innovative technologies and tools to improve species specificity and increase their feasibility on islands. *Biol. Conserv.* **185**, 47–58. <https://doi.org/10.1016/j.biocon.2014.10.016> (2015).
7. Gurnell, J. & Pepper, H. W. *The Control and Management of Grey Squirrel Populations in Britain* 407–436 (European Squirrel Initiative: European Squirrel Initiative, Suffolk, 2016).
8. Middleton, A. D. 38. the ecology of the American grey squirrel (*Sciurus carolinensis* Gmelin) in the British isles. In *Proceedings of the Zoological Society of London*, vol. 100, 809–843. <https://doi.org/10.1111/j.1096-3642.1930.tb01000.x> (Wiley Online Library, 1930).
9. Tompkins, D. M., Sainsbury, A. W., Nettleton, P., Buxton, D. & Gurnell, J. Parapoxvirus causes a deleterious disease in red squirrels associated with UK population declines. *Proc. R. Soc. Lond. Ser. B Biol. Sci.* **269**, 529–533 (2002).
10. England, N. *Lost life: England's Lost and Threatened Species* (Natural England, Peterborough, 2010).

11. Mountford, E. P., Peterken, G. F., Edwards, P. J. & Manners, J. G. Long-term change in growth, mortality and regeneration of trees in Denny wood, an old-growth wood-pasture in the new forest (UK). *Perspect. Plant Ecol. Evol. Syst.* **2**, 223–272. <https://doi.org/10.1098/rspb.2001.1897> (1999).
12. Hewson, C. & Fuller, R. *Impacts of Grey Squirrels on Woodland Birds: An Important Predator of Eggs and Young?* (British Trust for Ornithology, Thetford, 2003).
13. Williams, F. *et al.* The economic cost of invasive non-native species on great britain. *CABI Proj No VM10066* 1–99 (2010).
14. Grunwald, H. A. *et al.* Super-mendelian inheritance mediated by CRISPR-CAS9 in the female mouse germline. *Nature* **566**, 105–109. <https://doi.org/10.1038/s41586-019-0875-2> (2019).
15. Pfizner, C. *et al.* Progress toward zygotic and germline gene drives in mice. *CRISPR J.* **3**, 388–397 (2020).
16. Prowse, T. A. *et al.* Dodging silver bullets: Good CRISPR gene-drive design is critical for eradicating exotic vertebrates. *Proc. R. Soc. B Biol. Sci.* **284**, 20170799. <https://doi.org/10.1098/rspb.2017.0799> (2017).
17. McFarlane, G. R., Whitelaw, C. B. A. & Lillico, S. G. Crispr-based gene drives for pest control. *Trends Biotechnol.* **36**, 130–133. <https://doi.org/10.1016/j.tibtech.2017.10.001> (2018).
18. Burt, A. Site-specific selfish genes as tools for the control and genetic engineering of natural populations. *Proc. R. Soc. Lond. Ser. B Biol. Sci.* **270**, 921–928. <https://doi.org/10.1098/rspb.2002.2319> (2003).
19. Cong, L. *et al.* Multiplex genome engineering using CRISPR/CAS systems. *Science* **339**, 819–823. <https://doi.org/10.1126/science.1231143> (2013).
20. Galizi, R. *et al.* A CRISPR-CAS9 sex-ratio distortion system for genetic control. *Sci. Rep.* **6**, 31139. <https://doi.org/10.1038/srep31139> (2016).
21. Oberhofer, G., Ivy, T. & Hay, B. A. Cleave and rescue, a novel selfish genetic element and general strategy for gene drive. *Proc. Natl. Acad. Sci.* **116**, 6250–6259. <https://doi.org/10.1073/pnas.1816928116> (2019).
22. Champer, J. *et al.* A toxin-antidote CRISPR gene drive system for regional population modification. *Nat. Commun.* **11**, 1–10. <https://doi.org/10.1038/s41467-020-14960-3> (2020).
23. Simoni, A. *et al.* A male-biased sex-distorter gene drive for the human malaria vector anopheles gambiae. *Nat. Biotechnol.* **20**, 1–7 (2020).
24. Champer, J., Kim, I. K., Champer, S. E., Clark, A. G. & Messer, P. W. Performance analysis of novel toxin-antidote CRISPR gene drive systems. *BMC Biol.* **18**, 1–17. <https://doi.org/10.1186/s12915-020-0761-2> (2020).
25. Unckless, R. L., Clark, A. G. & Messer, P. W. Evolution of resistance against CRISPR/CAS9 gene drive. *Genetics* **205**, 827–841. <https://doi.org/10.1534/genetics.116.197285> (2017).
26. Champer, J. *et al.* Novel CRISPR/CAS9 gene drive constructs reveal insights into mechanisms of resistance allele formation and drive efficiency in genetically diverse populations. *PLoS Genet.* <https://doi.org/10.1371/journal.pgen.1006796> (2017).
27. Champer, J. *et al.* Reducing resistance allele formation in CRISPR gene drive. *Proc. Nat. Acad. Sci.* **115**, 5522–5527. <https://doi.org/10.1073/pnas.1720354115> (2018).
28. Champer, S. E. *et al.* Computational and experimental performance of CRISPR homing gene drive strategies with multiplexed grnas. *Sci. Adv.* **6**, eaaz0525. <https://doi.org/10.1126/sciadv.aaz0525> (2020).
29. Esvelt, K. M. & Gemmell, N. J. Conservation demands safe gene drive. *PLoS Biol.* <https://doi.org/10.1371/journal.pbio.2003850> (2017).
30. Noble, C. *et al.* Daisy-chain gene drives for the alteration of local populations. *Proc. Natl. Acad. Sci.* **116**, 8275–8282. <https://doi.org/10.1073/pnas.1716358116> (2019).
31. Min, J., Noble, C., Najjar, D. & Esvelt, K. Daisy quorum drives for the genetic restoration of wild populations. *BioRxiv* <https://doi.org/10.1101/115618> (2017).
32. Min, J., Noble, C., Najjar, D. & Esvelt, K. M. Daisyfield gene drive systems harness repeated genomic elements as a generational clock to limit spread. *BioRxiv* <https://doi.org/10.1101/104877> (2017).
33. Beaghton, A., Beaghton, P. J. & Burt, A. Vector control with driving y chromosomes: Modelling the evolution of resistance. *Malar. J.* **16**, 286. <https://doi.org/10.1186/s12936-017-1932-7> (2017).
34. Champer, J., Kim, I., Champer, S. E., Clark, A. G. & Messer, P. W. Suppression gene drive in continuous space can result in unstable persistence of both drive and wild-type alleles. *BioRxiv* <https://doi.org/10.1101/769810> (2019).
35. Robker, R. L., Akison, L. K. & Russell, D. L. Control of oocyte release by progesterone receptor-regulated gene expression. *Nucl. Receptor Signal.* **7**, nrs-07012. <https://doi.org/10.1621/nrs.07012> (2009).
36. Duarte, A. *et al.* Dosage-sensitive requirement for mouse dl4 in artery development. *Genes Dev.* **18**, 2474–2478. <https://doi.org/10.1101/gad.1239004> (2004).
37. Huerta-Cepas, J. *et al.* eggNOG 5.0: A hierarchical, functionally and phylogenetically annotated orthology resource based on 5090 organisms and 2502 viruses. *Nucl. Acids Res.* **47**, D309–D314. <https://doi.org/10.1093/nar/gky1085> (2019).
38. Champer, J., Buchman, A. & Akbari, O. S. Cheating evolution: Engineering gene drives to manipulate the fate of wild populations. *Nat. Rev. Genet.* **17**, 146. <https://doi.org/10.1038/nrg.2015.34> (2016).
39. Kyrou, K. *et al.* A CRISPR-CAS9 gene drive targeting doublesex causes complete population suppression in caged anopheles gambiae mosquitoes. *Nat. Biotechnol.* **36**, 1062–1066. <https://doi.org/10.1038/nbt.4245> (2018).
40. Dhole, S., Lloyd, A. L. & Gould, F. Tethered homing gene drives: A new design for spatially restricted population replacement and suppression. *Evol. Appl.* **12**, 1688–1702. <https://doi.org/10.1111/eva.12827> (2019).
41. Kandul, N. P., Liu, J., Bennett, J. B., Marshall, J. M. & Akbari, O. S. A home and rescue gene drive forces its inheritance stably persisting in populations. *BioRxiv* <https://doi.org/10.1101/2020.08.21.261610> (2020).
42. Champer, J. *et al.* A CRISPR homing gene drive targeting a haplolethal gene removes resistance alleles and successfully spreads through a cage population. *Proc. Natl. Acad. Sci.* **117**, 24377–24383 (2020).
43. Marshall, J. M. & Hay, B. A. Confinement of gene drive systems to local populations: A comparative analysis. *J. Theor. Biol.* **294**, 153–171. <https://doi.org/10.1016/j.jtbi.2011.10.032> (2012).
44. Edgington, M. P. & Alphey, L. S. Conditions for success of engineered underdominance gene drive systems. *J. Theor. Biol.* **430**, 128–140. <https://doi.org/10.1016/j.tree.2009.03.0160> (2017).
45. Gantz, V. M. *et al.* Highly efficient cas9-mediated gene drive for population modification of the malaria vector mosquito anopheles stephensi. *Proc. Natl. Acad. Sci.* **112**, E6736–E6743. <https://doi.org/10.1073/pnas.1521077112> (2015).
46. Lois, C., Hong, E. J., Pease, S., Brown, E. J. & Baltimore, D. Germline transmission and tissue-specific expression of transgenes delivered by lentiviral vectors. *Science* **295**, 868–872. <https://doi.org/10.1126/science.1067081> (2002).
47. Takahashi, G. *et al.* GONAD: Genome-editing via oviductal nucleic acids delivery system: A novel microinjection independent genome engineering method in mice. *Sci. Rep.* **5**, 11406. <https://doi.org/10.1038/srep11406> (2015).
48. Bradburd, G. S. & Ralph, P. L. Spatial population genetics: It's about time. *Annu. Rev. Ecol. Evol. Syst.* **50**, 427–449. <https://doi.org/10.1016/j.tree.2009.03.0163> (2019).
49. Rode, N. O., Estoup, A., Bourguet, D., Courtier-Ordogozo, V. & Débarre, F. Population management using gene drive: Molecular design, models of spread dynamics and assessment of ecological risks. *Conserv. Genet.* <https://doi.org/10.1007/s10592-019-01165-5> (2019).
50. Jones, H. E., White, A., Lurz, P. W., Boots, M. & Shuttleworth, C. M. *Mathematical Models of Grey Squirrel Invasion: A Case Study on Anglesey* 235–252 (European Squirrel Initiative: European Squirrel Initiative, Suffolk, 2016).

51. Gurnell, J. The effects of food availability and winter weather on the dynamics of a grey squirrel population in southern England. *J. Appl. Ecol.* **33**, 325–338. <https://doi.org/10.1016/j.tree.2009.03.0164> (1996).
52. Shorten, M. & Elton, C. Some aspects of the biology of the grey squirrel (*Sciurus carolinensis*) in Great Britain. *Proc. Zool. Soc. Lond.* **121**, 427–459. <https://doi.org/10.1111/j.1096-3642.1951.tb00804.x> (1951).
53. Barkalow, F. Jr., Hamilton, R. & Soots, R. Jr. The vital statistics of an unexploited gray squirrel population. *J. Wildl. Manag.* **34**, 489–500. <https://doi.org/10.1016/j.tree.2009.03.0166> (1970).
54. Thompson, D. The social system of the grey squirrel. *Behaviour* **64**, 305–328. <https://doi.org/10.1016/j.tree.2009.03.0167> (1978).
55. Thompson, D. Reproductive behavior of the grey squirrel. *Can. J. Zool.* **55**, 1176–1184. <https://doi.org/10.1016/j.tree.2009.03.0168> (1977).
56. Okubo, A., Maini, P. K., Williamson, M. H. & Murray, J. D. On the spatial spread of the grey squirrel in Britain. *Proc. R. Soc. Lond. B. Biol. Sci.* **238**, 113–125. <https://doi.org/10.1098/rspb.1989.0070> (1989).
57. Koprowski, J. L. *Sciurus carolinensis*. *Mammal. Spec.* <https://doi.org/10.2307/3504224> (1994).
58. Thompson, D. C. Regulation of a northern grey squirrel (*Sciurus carolinensis*) population. *Ecology* **59**, 708–715. <https://doi.org/10.2307/1938774> (1978).

## Acknowledgements

We thank Craig Shuttleworth for all his squirrel-related expertise and advice. C.B.A.W. acknowledges support from BBSRC ISP through BB/P013732/1 and BB/P013759/1. G.G. acknowledges support from the BBSRC to The Roslin Institute (BBS/E/D/30002275) and The University of Edinburgh's Data-Driven Innovation Chancellor's fellowship.

## Author contributions

C.B.A.W. and G.R.M. conceived the grey squirrel project. N.R.F. and G.R.M. conceived the novel gene drive. N.R.F. conducted the modelling with assistance from R.C.G., I.P., and G.G.. N.R.F. wrote the manuscript and all authors reviewed it.

## Competing interest

The authors declare no competing interests.

## Additional information

**Supplementary Information** The online version contains supplementary material available at <https://doi.org/10.1038/s41598-021-83239-4>.

**Correspondence** and requests for materials should be addressed to N.R.F.

**Reprints and permissions information** is available at [www.nature.com/reprints](http://www.nature.com/reprints).

**Publisher's note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2021